A sea spray aerosol flux parameterization encapsulating wave state

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Abstract. A new sea spray source function (SSSF), termed Oceanflux Sea Spray Aerosol or OSSA, was derived based on in-situ sea spray aerosol measurements along with meteorological/physical parameters. Submicron sea spray aerosol fluxes derived from particle number concentration measurements at the Mace Head coastal station, on the west coast of Ireland, were used together with open-ocean eddy correlation flux measurements from the Eastern Atlantic Sea Spray, Gas Flux, and Whitecap (SEASAW) project cruise. In the overlapping size range, the data for Mace Head and SEASAW were found to be in a good agreement, which allowed deriving the new SSSF from the combined dataset spanning the dry diameter range from 15 nm to 6 µm. The OSSA source function has been parameterized in terms of five lognormal modes and the Reynolds number instead of the more commonly used wind speed, thereby encapsulating important influences of wave height, wind history, friction velocity, and viscosity. This formulation accounts for the different flux relationships associated with rising and waning wind speeds since these are included in the Reynolds number. Furthermore, the Reynolds number incorporates the kinematic viscosity of water, thus the SSSF inherently includes dependences on sea surface temperature and salinity. The temperature dependence of the resulting SSSF is similar to that of other in-situ derived source functions and results in lower production fluxes for cold waters and enhanced fluxes from warm waters as compared with SSSF formulations that do not include temperature effects.

1 Introduction

Sea spray aerosol (SSA) is an important component of the aerosol population in the marine environment, and given that 70% of the Earth’s surface is covered by oceans, it contributes significantly to the global aerosol budget (Vignati et al., 2010). In addition, sea spray aerosol plays an important role in climate, with both direct (Bates et al., 2006; Mulcahy et al., 2008; Vaiyshya et al., 2011; Rap et al., 2013) and indirect radiative effects (O'Dowd et al., 1999; Andreae and Rosenfeld, 2008). Sea spray aerosol particles are formed at the sea surface mainly through breaking waves via bubble bursting (Blanchard, 1963) and, at elevated wind speeds, by direct tearing of wave crests (Monahan et al., 1986). When a wave breaks, air is entrained into the water and dispersed into a cloud of bubbles (Thorpe, 1992), which rise to the surface and burst to produce both film and jet drops. The near-surface wind speed, commonly measured and expressed at a reference height of 10 m, \( U_{10} \), is thought to be the dominant factor affecting sea spray aerosol production. However, different formulations of the size-dependent sea spray aerosol source functions (SSSF) in terms of only \( U_{10} \) vary widely for the same \( U_{10} \) (de Leeuw et al., 2011) and rising or waning winds produce different production fluxes (Callaghan et al., 2008; Goddijn-Murphy et al., 2011; Norris et al., 2012; Ovadnevaite et al., 2012). Considerable effort has been devoted to linking SSA production to more fundamentally relevant physical parameters, such as wind stress on the surface, or whitecap fraction, with the expectation that such
approaches might lead to better constraining the production flux than a formulation in terms of wind speed alone. However, at a given $U_{10}$, wind stress on the surface can vary by a factor of two (Drennan et al., 2005) and whitecap fraction by a factor of 10 or more (Lewis and Schwartz, 2004; Anguelova and Webster, 2006): this variation is likely due to variability in the wave field, surface properties, and the like. Factors that are expected to affect the SSA production flux are those affecting sea state, such as fetch (the distance an air mass has travelled over the water) and atmospheric stability (often parameterized as the air-sea temperature difference), which also affects vertical transport; seawater temperature and salinity; and the presence, amount, and nature of surface-active substances. The production of sea spray aerosol was recently reviewed by de Leeuw et al. (2011) who critically examined recent laboratory and field experimental results on sea spray aerosol production, on the enrichment in organic matter, and on the measurement and parameterization of whitecap fraction, and placed it in the context of previous understanding which was comprehensively reviewed by Lewis and Schwartz (2004). The study indicated that there is still considerable uncertainty in the production of sea spray aerosol and suggested that existing submicron number flux parameterizations appear to over-predict boundary layer number concentrations compared to what is actually measured.

In this study we derive a new sea spray aerosol source function, termed Oceanflux Sea Spray Aerosol or OSSA, which covers the dry particle size range from 15 nm to 6 µm in diameter and uses a direct fit of the size-dependent flux observations to a wave breaking parameter (Reynolds number). This approach combines the effects of wind history and wave state along with the sea surface temperature and salinity in one parameter.

2 Approach

2.1 SSA flux formulation

Sea spray aerosol generation is directly related to wind-induced breaking of waves, which entrains air into the ocean surface layer. The entrained air is detrained as bubbles, which rise to the surface where they burst and produce film and jet droplets (Blanchard, 1986). The surface manifestation of a bubble plume is referred to as a whitecap and the fraction of the ocean surface covered by whitecaps, i.e. the whitecap fraction, is often used as the basis for the formulation of a sea spray aerosol source function (Lewis and Schwartz, 2004). In this so-called whitecap method, the whitecap fraction is parameterized in terms of a forcing parameter (e.g. 10 m wind speed or Reynolds number), and the size-dependent droplet production per unit whitecap. Hence, only the magnitude of the flux is allowed to vary with the forcing parameter (de Leeuw et al., 2011). This assumption is an oversimplification of the real environment, where the shape of the droplet size distribution could and most certainly does depend on the forcing parameter. Therefore, in this study, we incorporate the effect of the environmental forcing on both the magnitude and the shape of the SSSF. In other words, a direct relationship is assumed between the sea spray aerosol flux and the forcing parameter. Several processes would affect the bubble bursting and thus resulting aerosol or droplet size distributions; therefore, a size dependent flux parameterization is assumed to include several modes, which, at the surface, should follow a normal distribution. On the other hand, after initial formation the droplets are dispersed in the atmosphere (multiple dilutions into volume), which transforms the distribution into a lognormal distribution (Limpert et al., 2001).

Since the current large-scale aerosol models cannot resolve this micro-scale aerosol dynamics, we directly assign a lognormal distribution to the source flux modes, which should thus be considered effective fluxes. Each mode is then linked to the forcing parameter, because the magnitudes of separate modes could depend on it in a different way.

2.2 Forcing factors

Wind waves are generated by transferring wind energy to the water surface through friction. Continuous wind stress increases the wave height until the wave breaks which in turn results in energy dissipation. Wave age and sea state will usually depend upon wind history, e.g. periods of decreasing wind speed would correspond to more developed seas with a relatively large wave age and periods of increasing wind speed should be broadly analogous to less developed seas with a relatively small wave age (Callaghan et al., 2008). Based on consideration of the energy flux from wind to waves, Wu (1979) proposed that wave breaking, and therefore whitecap fraction, should be proportional to $u_*^3$ or $U^{3.75}$ where $u_*$ is the friction velocity. At about the same time, Monahan and Muircheartaigh (1980) proposed their $U^{3.10}$ power law derived from a large amount of whitecap observation datasets. However, the wave state depends not only on the actual wind speed but also on wind history (fetch, increasing or decreasing winds), leading to developing or well-developed waves, with different wave steepness. Therefore, based on physical considerations of wave breaking, Zhao and Toba (2001) proposed the use of a dimensionless breaking wave parameter ($R_b$); however, the typical length scale to construct $R_b$ was ambiguous and therefore was replaced by $H_w$, which in turn led to a Reynolds number ($Re_{H_w}$) expression. The latter includes the friction velocity, the height of wind waves, and the viscosity of the air. Zhao and Toba (2001) further proposed to replace the viscosity of air with the viscosity of water which is conceptually more relevant, and which was later reinforced by Woolf (2005). The resulting expression for the Reynolds number $Re_{H_w}$ is presented below:
\[ Re_{H_w} = \frac{u_w H_w}{v_w} \]  

Here \( H_w \) is the significant wave height and \( v_w \) is the viscosity of water. The viscosity of sea water depends on temperature and salinity (Sharqawy et al., 2010), the effects of which are thus implicitly included in a sea spray aerosol source function formulated in terms of \( Re_{H_w} \). \( Re_{H_w} \) increases with increasing temperature (decreasing viscosity) and vice versa. The use of either the breaking wave parameter or the Reynolds number brings results from several laboratory experiments and in-situ observations for whitecap fraction together (Zhao and Toba, 2001), unlike the use of parameters like (non-dimensional) fetch or wave age. The good correlation between whitecap fraction and breaking wave parameter was further demonstrated by Goddijn-Murphy et al. (2011).

### 2.3 SSSF expression

Considering the reasons presented above, the OSSA SSSF is expressed as a combination of \( n \) lognormally distributed modes for different droplet sizes, each of which with distinct dependency on the forcing parameter (Eq. 2). The Reynolds number was selected as the forcing parameter because of the advantages presented in Sect. 2.2.

\[
\frac{dF}{d \log D} = \sum_{i=1}^{n} F_i \left( Re_{H_w} \right) \frac{1}{\sqrt{2\pi} \ln \sigma_i} \exp\left(-\frac{1}{2} \left( \frac{\ln(D/CMD)}{\ln \sigma_i} \right)^2 \right) \]  

Here \( \frac{dF}{d \log D} \) is the size dependent SSA production flux, \( i \) – mode number, \( F_i (Re_{H_w}) \) – the flux for mode \( i \), \( D \) – dry particle diameter, \( \sigma \) – geometrical standard deviation, CMD – count median diameter.

Ambient observations, described below, were used to derive the specific \( F_i (Re_{H_w}) \) relationships and the appropriate parameters defining the lognormal modes.

### 3 Description of the data

The source function was derived from the combination of three datasets: sub-micron particle size distributions measured at Mace Head, coarse mode SSA fluxes derived from open-ocean eddy-covariance measurements during the SEASAW cruise (Norris et al., 2012) and wave parameters obtained from the European Centre for Medium-Range Weather Forecasts (ECMWF) model. Mace Head and SEASAW data did not cover the same time and region, but both datasets are representative for the North East Atlantic during low biological productivity and cover complementary size ranges.

#### 3.1 Submicron particle observations

Submicron aerosol particle size distributions were used, which were measured at the Mace Head Atmospheric Research Station in November 2010. Mace Head is located on the west coast of Ireland facing the North East Atlantic. Station details are described in O’Connor et al. (2008). All aerosol instruments are located in the shore laboratory at about 100 m from the coastline. They are connected to the lamar flow community air sampling system, which is constructed from a 100 mm diameter stainless-steel pipe with the main inlet at 10 m above ground level. The performance of this inlet is described in Kleefeld et al. (2002). Details on the representativeness of marine aerosol measured at Mace Head for open-ocean conditions can be found in the recent study by O’Dowd et al. (2013).

Aerosol size distributions and number concentrations were measured using a scanning mobility particle sizer (SMPS) system. The system comprised of a differential mobility analyser (DMA, TSI model 3071), a condensation particle counter (TSI model 3010), and an aerosol neutralizer (TSI 3077). Before their sizes were measured, the particles were dried to a relative humidity below 40%.

The wind speed and direction were measured on the 10 m tower by a Vector Instruments wind monitor (model W200P/A100L).

Aerosol chemical composition was measured using a High Resolution Time of Flight Aerosol Mass Spectrometer (HR-ToF-AMS or AMS) (DeCarlo et al., 2006) which was routinely calibrated following the methods described by Jimenez et al. (2003) and Allan et al. (2003). The HR-ToF-AMS measurements were performed with a time resolution of 5 min and a vaporizer temperature of \( \sim 650^\circ \text{C} \). The composition dependent collection efficiency (CE) (Middlebrook et al., 2012) was applied for the measurement periods discussed in this study. Sea salt concentrations were measured following the method described in Ovadnevaite et al. (2012).

Data obtained during the occurrence of a plume with elevated sea salt concentrations over the North East Atlantic, see Ovadnevaite et al. (2012) for details, were used to derive the submicron SSSF. The plume was detected as the wind direction backed northerly into the clean sector at Mace Head (between 190\(^\circ\)-300\(^\circ\)) and the wind speed increased to a peak value of 26 m s\(^{-1}\). Sea salt plumes measured by the HR-ToF-AMS coincided with an increase in aerosol hygroscopicity from a typical sulphate hygroscopic growth factor (GF) of 1.6 to a GF of 2.2, which is characteristic of pure sea salt particles. As the measurements were undertaken during the winter when biological activity was low, all other chemical compounds approached very low background “winter” concentrations (e.g. non-sea salt sulphate mass < 10 ng m\(^{-3}\); organic mass < 60 ng m\(^{-3}\); black carbon mass < 10 ng m\(^{-3}\); nitrate mass < 17 ng m\(^{-3}\) and ammonium mass below the detection limit of 38 ng m\(^{-3}\)).

#### 3.2 Super-micron particle observations

Since Mace Head data covers only the submicron part of the SSSF, fluxes of larger sea spray aerosol particles measured during the SEASAW campaign (described and analysed in
detail in Norris et al., 2012) were used to complement the Mace Head derived SSSF. The SEASAW open-ocean flux dataset consists of 111 valid measurements obtained during a cruise in the Eastern North Atlantic Ocean, 21 March–12 April 2007. Wind speed conditions ranged from 3 to 18 m s\(^{-1}\), and the ship was actively relocated to areas with high wind speed conditions. The biological activity, estimated from satellite data and aerosol volatility measurements, was low. Data were obtained with the Compact Lightweight Aerosol Spectrometer Probe (CLASP) (Hill et al., 2008), measuring aerosol particle concentrations in 16 size channels covering a range of 0.17–9.5 µm radius at ambient relative humidity. Sea spray aerosol fluxes were measured using the eddy covariance technique, with sampling records of 28 min.

Flux estimates were corrected for the relative humidity flux, and results were harmonized to represent fluxes for particle radii at 80% relative humidity. In the literature, it is common to use either the dry particle diameter (\(D_{dry}\)) or the radius at 80% RH (\(R_{80}\)): in fact these two conventions come down to the same numerical value as for sea salt particles a wet radius at 80% RH equals the dry diameter. When using these data to derive an SSSF, there are some limitations which should be mentioned. Firstly, the scatter in the data was considerable (order of magnitude), which is inherent to the measurement technique. Secondly, a net flux was measured, which includes both the production flux and the deposition. Norris et al. (2012) indicate that the difference between the production flux and the net flux is important for super-micron particles larger than about \(R_{80} = 5 \mu m\), however, this difference becomes insignificant for particles smaller than 1 µm.

### 3.3 Wave data

Wave data from the ECMWF wave model (WAM) were used in this study to derive the Reynolds number from wave characteristics. Data were globally available on 6h basis at a 0.5 x 0.5° longitude-latitude resolution. The ECMWF Re-Analysis (ERA-interim) product was used to get a consistent dataset over several years. We have used the wind velocity, mean drag coefficient, and significant height of wind waves. WAM assimilates wave height data derived from satellite altimetry data (Abdalla et al., 2010).

### 4 Derivation

#### 4.1 The expression of the Reynolds number

The \(Re_{Hw}\) expression used in this paper (Eq. 3) is obtained by replacing \(u_\ast\) in Eq. (1) with \(u_\ast = C_1^{1/2} U_{10}\). To avoid an influence from a swell component, only the wind wave part of the wave height, as provided by ECMWF, was used in the SSSF derivation.

\[
Re_{Hw} = C_1^{1/2} U_{10} H_w / \nu_w
\]

This replacement of \(u_\ast\) only holds for neutral conditions. However, this approach is deployed in WAM, which output (\(H_w, C_d\) and \(U_{10}\)) is used in this study. In addition, it takes into account the calculated wave state: the WAM model is coupled to the ECMWF atmosphere model with a correction for the Charnock parameter based on the wave stress in the feedback of the wave model to the atmosphere model (IFS, 2011). Foreman and Emeis (2010) proposed a new relationship between \(u_\ast\) and \(U_{10}\), based on a fit to wind and wave observations, later modified by Andreas et al. (2012), which could be an alternative to the formulation that we have used, but then the coupling to wave state and the internal consistency in our dataset would be lost. More importantly, their average relationships cannot resolve the subtleties of the effect of rising or waning winds or fetch on wave state, which is included in WAM. It is interesting to note that in the parameterizations of Foreman and Emeis (2010) and Andreas et al. (2012), a distinction is made in fit parameters between wind speeds smaller than or larger than 8–9 m s\(^{-1}\), related to the transition to rough flow, with impact on the drag. Such distinction between different wind speed regimes has also been made in studies on whitecap parameterization (Callaghan et al., 2008; Godijn-Murphy et al., 2011); however without reference to this transition to rough flow. The direct use of \(u_\ast\) in the parameterization has the advantage that the need to separate into different wind speed regimes disappears, provided that the relationship between \(u_\ast\), wind speed and drag is realistic. The fact that we did not see any transition in our source function for different wind speed regimes, as opposed to what is usually seen in wind speed-only source functions, supports the idea that our approach captures the main features.

Moreover, using the SSSF based on wind speed only, the mass flux for waning winds significantly exceeds the flux for rising winds (Ovadnevaite et al., 2012) and the flux-wind speed dependencies for these two different wind conditions is clearly separated (Fig. 1, right panel, and Ovadnevaite et al., 2012). For the new SSSF, parameterized in terms of \(Re_{Hw}\), the difference between rising and waning winds disappeared. This not only reduced the scatter (\(R^2\) improved from 0.95 to 0.98 and chi square reduced from 16.4 to 5.8) but the relationships for rising or waning winds started to inter-cross (Fig. 1, left panel), which also indicated that remaining data point scatter was due to a data measurement or derivation uncertainty rather than a real physical effect coming from the wave state.

#### 4.2 Submicron SSA flux calculations

Submicron SSA fluxes were derived from the particle number concentration measurements using the following assumptions. Submicron aerosol particles were expected to
Fig. 1. Production Flux dependence on the Reynolds number, $R_{Hw}$, (left panel) and wind speed (right panel). For the data points presented here, the wind speed was initially increasing then levelling off and eventually decreasing, therefore, intercrossing lines for the relationship on the left panel – Flux vs. Reynolds number – indicate that there is no separation between the two regimes (increasing and decreasing wind speed), however, Flux vs. Wind speed (right panel) indicates distinctly different relationships for the different regimes, especially at higher winds.

be uniformly mixed in the marine boundary layer (Lewis and Schwartz, 2004), thus an effective SSA production flux, $F_{\text{eff}}$, was estimated from the SSA number concentration ($N$) divided by a filling time ($\tau$) and multiplied by the marine boundary layer height ($H_{\text{MBL}}$):

$$F_{\text{eff}} = \frac{N \times H_{\text{MBL}}}{\tau} \tag{4}$$

The atmospheric boundary layer height was derived from ground-based LIDAR measurements using the Temporal Height Tracking (THT) algorithm (Haefelin et al., 2012; Milroy et al., 2012). The marine boundary layer typically consists of two layers, with a surface mixed layer (SML) and a decoupled residual or convective layer (DRCL) above which is the free troposphere (Milroy et al., 2012). SML values were used in the flux calculations and were observed to vary between 720 and 1290 m above ground level over the plume duration period. SML height derived over Mace Head was comparable to the relatively steady mixing layer depth over the ocean observed along the air mass back trajectories. Nonetheless, remaining differences between the SML and the marine boundary layer height contributed to the SSSF uncertainties.

The filling time $\tau$ was assumed to be approximately 1.5–2 days, as discussed in Ovdanvaite et al. (2012) who used a similar method to derive the submicron sea spray aerosol mass flux.

The local wind speed at Mace Head was representative of open ocean conditions further upwind, obtained from the National Oceanic and Atmospheric Administration (NOAA) Air Resources Laboratory, as illustrated in Fig. 2a. Moreover, Fig. 2c–d indicate the good agreement between wind speeds measured at Mace Head and the wind speeds modelled by ECMWF for the area upwind to Mace Head (Fig. 2b) which, together with ECMWF model data for significant wave heights and drag coefficients, was used to calculate $R_{Hw}$. Since ECMWF data were provided with a resolution of only 6 hours, too coarse for application to the relatively short event studied here, they were interpolated to a resolution of 10 min. Modeled, interpolated and measured data are presented in Fig. 2c, d.

Although the background mass concentration (as measured at $\sim$4 m s$^{-1}$ wind speed) was insignificant (Ovdanvaite et al., 2012), the background number size distribution was subtracted from the number distributions measured at higher wind speeds.

Using deposition velocities provided in (Hoppel et al., 2002), the dry deposition was found to be negligible for the submicron particles, except for the very small nano-metric size particles or for the super-micron ones; however, the error introduced due to deposition at these sizes would still be within the uncertainties provided for the OSSA source function. During the period of Mace Head observations used here, there were practically no clouds or precipitation and therefore the contribution of wet-deposition and coalescence removal processes to the total flux were negligible.

4.3 OSSA-SSSF

The size dependent SSA production flux ($dF/d\log D$) was obtained by combining a submicron aerosol flux, derived from $dN/d\log D$ data (measured by SMPS) and calculating $F$ from $N$ using Eq. (3), with the measured super-micron $dF/d\log D$ data and averaged over 1 m s$^{-1}$ wind speed bins. An average Reynolds number was also calculated for each of these bins. Combined $dF/d\log D$ data points are presented in Fig. 3 for different wind speeds. In the overlapping size range (at $\sim$300 nm) the two datasets show very good agreement, except for the measurements at the lowest and the highest wind speeds, although these were still within
the uncertainty limits described above. For every wind speed bin, five lognormal size distributions were fitted to the resulting $dF/d\log D$ distribution (Fig. 4). This resulted in a five-modal SSA flux formulation in terms of the Reynolds number $Re_{H_w}$ with different coefficients for each mode (Eq. 2 with $n = 5$).

All parameters used in Eq. (2) are listed in Table 1, including the SSA production flux relationship with $Re_{H_w}$ through the $F_i(Re_{H_w})$.

The dependence of $F_i$ on $Re_{H_w}$ is shown in Fig. 4b together with the measurement data. The $Re_{H_w}$ dependence is distinctly different for each mode, which confirmed the assumption, raised in the “Approach” section, that the forcing parameter affects the production of particles of different sizes.

Table 1. Lognormal parameters for the SSSF parameterization. For each mode, a geometric standard deviation ($\sigma_i$), count-median diameter (CMD$_i$), and the total number flux ($F_i$) as a function of Reynolds number ($Re_{H_w}$) are given.

<table>
<thead>
<tr>
<th>$i$</th>
<th>$\sigma_i$</th>
<th>CMD$_i$</th>
<th>$F_i(Re_{H_w})$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1.37</td>
<td>0.018</td>
<td>$104.5(Re_{H_w}-1 \times 10^5)^{0.556}$</td>
</tr>
<tr>
<td>2</td>
<td>1.5</td>
<td>0.041</td>
<td>$0.0442(Re_{H_w}-1 \times 10^5)^{1.08}$</td>
</tr>
<tr>
<td>3</td>
<td>1.42</td>
<td>0.09</td>
<td>$149.6(Re_{H_w}-1 \times 10^5)^{0.545}$</td>
</tr>
<tr>
<td>4</td>
<td>1.53</td>
<td>0.23</td>
<td>$2.96(Re_{H_w}-1 \times 10^5)^{0.79}$</td>
</tr>
<tr>
<td>5</td>
<td>1.85</td>
<td>0.83</td>
<td>$0.51(Re_{H_w}-2 \times 10^5)^{0.87}$</td>
</tr>
</tbody>
</table>
sizes differently. The study by Monahan et al. (1986) indicated various wind speeds for the onset of the production by different mechanisms. Therefore, separate intercepts for submicron and super-micron aerosol particles ($Re_{H_w}$ at $1 \times 10^5$ and $2 \times 10^5$ respectively in Table 1) were anticipated. The $Re_{H_w}$ threshold for submicron particles is consistent with results from the study by Callaghan et al. (2008), which suggested a wind speed threshold for the onset of wave breaking at $\sim 3.7 \text{ m s}^{-1}$, corresponding to $Re_{H_w} \sim 1 \times 10^5$ at given wave height and sea surface temperature (SST) conditions ($H_s = 1.23$, SST $= 9^\circ\text{C}$).

### 4.4 Error propagation

The SSSF uncertainty was evaluated by applying error propagation rules (Taylor, 1997). The submicron aerosol flux was calculated from the number concentration ($N$), the boundary layer height ($H_{MBL}$) and the filling time ($\tau$) (see Eq. 3), therefore, the resulting uncertainty was equal to $\sqrt{\Delta N^2 + \Delta H_{MBL}^2 + \Delta \tau^2}$. The SMPS measurement uncertainty $\Delta N \sim 10–20\%$, $\Delta D \sim 5\%$ (Wiedensohler et al., 2012), $\Delta H_{MBL} \sim 15–20\%$ (Milroy et al., 2012) and $\Delta \tau \sim 50\%$ (Ovadnevaite et al., 2012), therefore, the resulting submicron SSSF uncertainty is in the range of $55–60\%$

Since the derived Reynolds number was used instead of direct $U_{10}$ measurements, the uncertainty due to $Re_{H_w}$ calculations (see Eq. 2) was evaluated accordingly: $\sqrt{\Delta C_d^2 + \Delta U_{10}^2 + \Delta H_{MBL}^2}$; the uncertainty in $\Delta H_s$ is $\sim 2–5\%$. $\Delta U_{10} \sim 10\%$, $\Delta C_d \sim 15\%$ (Janssen et al., 2007; Abdalla et al., 2011), which resulted in an overall $Re_{H_w}$ uncertainty of $\sim 20\%$.

Super-micron aerosol measurement and eddy covariance method uncertainties were presented in the study by Norris et al. (2012), Sects. 2.2 and 2.3.

### 5 Results

Measured and parameterized fluxes for two different values of $Re_{H_w}$ are presented in Fig. 5, which shows $dF/d\log D$ vs. $D$ together with the calculated uncertainties. Figure 5 shows that the parameterized fluxes represent the observations well within the uncertainty range, which suggests that the main driving processes were captured correctly.

#### 5.1 SSSF intercomparison

The OSSA-SSSF has been evaluated by comparing the resulting sea spray aerosol mass with independent HR-ToF-AMS measurements at Mace Head, which were not used in the derivation of the OSSA-SSSF. However, the data can only be used to validate the submicron part of the spectrum. In order to achieve this, the OSSA number flux was simulated for a particle size range directly comparable to the HR-ToF-AMS measurement size range ($D_{MD} = 0.03–0.58 \mu m$ or vacuum aerodynamic diameter, as measured in the HR-ToF-AMS, $D_{va} = 0.05–1 \mu m$), converted to a corresponding mass flux and integrated over the size range. This was done for a range of wind speeds and the results are shown in Fig. 6, together with HR-ToF-AMS derived mass fluxes (Ovadnevaite et al., 2012) and the respective uncertainties. The production flux derived using the OSSA-SSSF is within the uncertainty ranges of the HR-ToF-AMS mass fluxes although it overestimates the mass concentration from $0\%$ (at $5 \text{ m s}^{-1}$ wind speed) to $80\%$ (at $20 \text{ m s}^{-1}$ wind speed). However, the agreement between the OSSA-SSSF and the HR-ToF-AMS derived SSA mass fluxes is significantly better than that between HR-ToF-AMS and other SSSFs presented in Ovadnevaite et al. (2012). The largest discrepancy between the SSA mass fluxes obtained with the OSSA-SSSF and measured with the HR-ToF-AMS is due to the large uncertainty in super-micron aerosol distributions (see SEASAW description in Sect. 2) since SSA particles in this size range have the highest contribution to the SSA mass. As shown in Fig. 4, particles in mode 5 contribute significantly to the submicron aerosol number concentration (see $F_5$ in Fig. 4a) and therefore also to the resulting mass flux together with the inherent uncertainties.

In addition, the variation in the HR-ToF-AMS aerodynamic lens cut-off could have had an effect on the mass concentrations measured by the HR-ToF-AMS. The typical HR-ToF-AMS aerodynamic lens cut-off of $1 \mu m$ (50\% transmission efficiency for the $D_{va}$ of $1 \mu m$ or mobility diameter, $D_m$, of $0.58 \mu m$) was applied to the number flux derived from this study in order to compare different methods. However, the best agreement between the masses derived from
the OSSA-SSSF and measured by the HR-ToF-AMS was achieved when an upper cut-off of 0.9 µm is selected.

A comparison of the OSSA sea spray aerosol source function (OSSA-SSSF) and other commonly used or recently (last decade) developed source functions at 8 m s\(^{-1}\) wind speed (de Leeuw et al., 2011) is presented in Fig. 7. This wind speed was selected as the most common one in the real ambient environment (Rinaldi et al., 2013). Since the OSSA-SSSF depends on the Reynolds number instead of the wind speed directly, \(Re_{H_w}\) was calculated using Eq. (2) for a wind speed of 8 m s\(^{-1}\) and other parameters relevant to the conditions for which this source function was derived (\(C_d = 2.15 \times 10^{-3}\), \(H_s = 1.23\), \(\nu_w = 1.34 \times 10^{-6}\)).

The OSSA-SSSF is on the lower side as compared to other parameterizations. As noted by de Leeuw et al. (2011) and Ovadnevaite et al. (2012), source functions based on Monahan’s whitecap parameterization tend to result in higher atmospheric submicron number/mass concentrations than those typically measured. A recent European Monitoring and Evaluation Programme (EMEP) unified model intercomparison study (Tsyro et al., 2011) shows that model calculations using the Mårtensson et al. (2003) and Gong (2003)
SSSF overestimate atmospheric concentrations of Na by 8% to 46% in comparison with EMEP observations. The higher overestimation was observed for the winter season, which is consistent with the results presented in Ovdanevaite et al. (2012): the stronger flux-wind speed relationship of Mårtensson et al. (2003) or Gong (2003) would result in higher overestimation of sea salt mass concentrations during the high wind speed periods usually occurring in winter.

5.2 An estimate of the global production flux

Although validation of the OSSA-SSSF on a global scale was performed elsewhere (Partanen et al., 2014), it has been used for the preliminary calculation of the annual mean production flux for the year 2006. This was achieved with a simple modelling tool, developed at Netherlands Organisation for Applied Scientific Research: TNO, which calculates the fluxes based on the prescribed parameterization, and uses ECMWF meteorological and wave data as an input (Fig. 8).

Mass and number fluxes were calculated for particles with dry diameters < 1 µm. 3 h sea surface temperature and $U_{10}$ from ECMWF Integrated Forecasting System (IFS) analysis, 6 h $C_d$, significant height of wind waves from WAM ERA-Interim and sea surface temperature (SST) dependent viscosity, assuming a constant salinity of 35‰, were used to calculate $Re_{H_s}$ using Eq. (2). The Reynolds number was then used in the parameterization (2) with the parameters of Table 1. It is worth noticing that appropriate salinity and thus viscosity values should be used (but not applied in this study) when the source function is applied to brackish waters such as the Baltic Sea (Mårtensson et al., 2003; Manders et al., 2010; Sofiev et al., 2011).

As expected, Fig. 9 shows higher production fluxes for the regions with stronger winds. In addition, sea surface temperature effects, incorporated in $Re_{H_s}$ through the viscosity, enhance the production in the warm waters on both sides of the equator, and reduce it for the colder water regions with respect to an exercise in which the OSSA-SSSF was used with constant viscosity (Fig. 10).

Global instantaneous fluxes derived from the meteorological and wave data are presented in Fig. 11, which shows a comparison of these fluxes with those derived with the same modelling tool but using the Gong (2003) and Jaeglé et al. (2011) source functions. These two source functions are similar with the difference that Jaeglé et al. (2011) incorporate SST effects based on a comparison between model computations and MODIS observations of the aerosol optical depth (AOD). The comparisons were made for instantaneous global data (2 January 2006, 00:00 UTC) for number fluxes of particles with a dry diameter of $0.07 < D_{dry} < 0.15$ µm (Aitken mode) or $0.07 < D_{dry} < 1$ µm (submicron particles) and for the mass flux of particles smaller than 1 µm. Fluxes computed using the Gong SSSF are presented on both left and right panels in black dots, while the fluxes computed using the OSSA-SSSF are presented in coloured circles on the left panel and those using the Jaeglé SSSF in coloured circles on the right panel. The differences in the fluxes due to the SST effect, as compared to the Gong SSSF, are obvious for both the OSSA and the Jaeglé et al. (2011) SSSF. Higher fluxes are produced at higher SST (red colours in the Fig. 11); while at lower temperatures (blue colours) the fluxes are lower.

In addition to the spread due to SST, the OSSA-SSSF also accounts for wave state, which reduces the effect of...
Fig. 8. Annual mean values of meteorological/oceanographic fields (ECMWF) that were used for the calculation of the sea spray aerosol fluxes for 2006.

temperature on the fluxes and brings some of the low temperature points closer to the Gong-derived fluxes. We postulate that this is due to on average larger values of the wave height and drag coefficient in the lower temperature regions (Fig. 8). As an example, the low flux values calculated using the Jaeglé SSSF at high wind speeds and at low SST (Fig. 11a, right panel) are not observed in the fluxes calculated with the OSSA-SSSF (Fig. 11a, left panel). The submicron number flux resulting from our function is quite evenly distributed around Gong’s function for all wind speeds (Fig. 11a, left panel). However, the number fluxes for the Aitken mode particles, important for cloud formation, are lower than those obtained using Gong, especially at higher wind speeds (Fig. 11b, left panel). In addition, the OSSA-SSSF derived submicron mass fluxes are considerably lower than those derived using Gong’s SSSF, at all temperatures except for the highest ones (Fig. 11c, left panel).

In order to compare the temperature dependences of the OSSA and Jaeglé et al. (2011) SSSFs and considering that the Jaeglé et al. (2011) SSSF does not have a dependency on the wind history, averaged constant values for $C_d$ and $H_s$ were used in the flux calculations applied for the comparison. Although different in origin (through the viscosity in the OSSA-SSSF and through the adjustment to the in-situ measurements in the Jaeglé SSSF), the variations of the SSSF with SST are similar in the sense that for both SSSFs the production increases with SST and the values for the lowest and highest temperatures are similar (Fig. 12). However, the shapes of the relationships are somewhat different. Where the OSSA-SSSF increases monotonously with SST, the Jaeglé
Fig. 9. Annual mean production flux of SSA computed using the OSSA-SSSF together with ECMWF meteorological and wave information: (a) Number flux of particles smaller than 1 µm; (b) Mass flux of particles smaller than 1 µm; (c) Mass flux of particles between 1–2.5 µm.

et al. (2011) SSSF is much lower in the mid-temperature regime, but rapidly increases at higher temperatures. At this stage, it is difficult to say which dependency is more appropriate as the OSSA-SSSF SST dependency derives from first principles but is not (yet) compared to observational data representing different SST, while Jaeglé’s comes from the adjustment to AOD measurements; however, AOD measurements are not a good proxy for the direct particle flux since AOD is determined by both production and subsequent processes in the atmosphere, i.e. transport, removal, and transformation due to chemical and physical processes which affect particle concentrations, size distributions, and optical properties. Also, AOD over ocean may be affected by aerosol particles of non-marine origin.

6 Discussion

Introducing the Reynolds number into the OSSA-SSSF brought about improvements as regards the ambiguity in the effect of wind speed and the sensitivity to other environmental parameters. First of all, the scatter at a given wind speed arising from wind history (fetch, rising or waning winds) was reduced by including the wave state in the forcing parameter. Secondly, the Reynolds number encapsulated the SST effects through the viscosity of sea water which depends on its temperature and salinity, however, effect of SST has not been validated yet. Several studies have shown the impact of SST on the SSSF (Mårtensson et al., 2003; Sellegri et al., 2006; Jaeglé et al., 2011; Zábori et al., 2012), but the results from the laboratory experiments were somewhat contradictory or at least not consistent. Mårtensson et al. (2003) showed a decrease in particle number concentration with increasing temperature for particles smaller than 0.35 µm, but an increase for larger particles, while Sellegri et al. (2006) indicated a different threshold at ~0.07 µm; therefore, particles with a diameter of ~0.1 µm would decrease in number concentration for one SSSF, but increase for another one. At the same time, the recent study of Zábori et al. (2012) showed a decrease in number concentration for all
particles smaller than 2.5 µm with increasing temperature. In contrast, ambient mass measurement studies (Jaegle et al., 2011 and references therein) indicated an increase in particle mass with increasing temperature. This mass-temperature dependency is similar to the temperature dependency captured in the OSSA-SSSF, where SST effects are included through the kinematic viscosity of seawater. The latter was easy to implement into the SSSF and derives from first physical principles, because the terminal velocity of a rising bubble is inversely proportional to the kinematic viscosity of the surrounding fluid (Hinds, 1982). Thus, bubbles in warmer waters will rise more quickly to the surface than in colder waters (e.g. bubbles of 0.04 mm reach terminal rise velocities of 0.28 cm s$^{-1}$ and 0.53 cm s$^{-1}$ at 0 and 20°C, respectively, Lewis and Schwartz, 2004), which would increase the number of smaller bubbles reaching the surface, and thus increase the production of SSA particles (Jaegle et al., 2011). Also Anguelova and Webster (2006) indicated that there is an
In particular, the representativeness of small scale laboratory parameterizations based on laboratory experiments are facing, and SEASAW; therefore, it lacks many problems which parameterizations developed by Rinaldi et al. (2013) in a way it was used before in several other studies (O’Dowd et al., 2008; Vignati et al., 2010; Gantt et al., 2011). There is also evidence that micro-organisms affect the viscosity of sea water (Seuront et al., 2010) so that biological activity may be taken into account via the viscosity, like the effect of temperature and salinity; however, further studies and parameterizations are required on this topic in order to separate the different effects and relate the viscosity to observables like chlorophyll-a concentrations.

7 Conclusions

The OSSA sea spray aerosol source function derived in this study covers particle sizes with dry diameters between 15 nm up to 6 µm and encapsulates wave history, salinity, and temperature effects through using the Reynolds number as forcing parameter instead of the wind speed. For the first time, this source function shows and accounts for the distinct forcing effects on the different particle size ranges, resulting in different flux distributions for a particular $Re_{H_w}$. The resulting SSSF provides fluxes which are on the lower side of those calculated using many other source functions developed in the last decade, yet the use of the OSSA-SSSF results in particle number and mass concentrations closer to the ones measured in real ambient conditions. Better agreement with the measurements allows producing more accurate particle number concentrations and size distributions, which in turn results in a better description of cloud condensation nucleus distributions. The latter is very important in order to reduce the uncertainty in modelling indirect effects on the earth radiative balance arising from primary production of SSA. In addition to climate effects, a correct particulate mass assignment to the natural sources, in this case primary marine sea spray aerosol production, is very important as regards air quality and source apportionment studies.

Another important advantage of the OSSA-SSSF is that it was derived from in-situ observations, from Mace Head and SEASAW; therefore, it lacks many problems which parameterizations based on laboratory experiments are facing, in particular, the representativeness of small scale laboratory experiments for oceanic conditions. At the same time, coastal stations possess the risk of the effects of the surf zone on direct measurements of SSA fluxes. In this study, the SSA fluxes were derived from concentration measurements which are not affected by the coastal effects (Ceburnis et al., 2008). In addition, the specific topography of Mace Head minimises the influence of the surf zone as well (Rinaldi et al., 2009). Moreover, the fact that the two independent and methodologically different datasets used in this study (Mace Head and SEASAW) were so similar in the overlapping size range provided us further confidence in the results.

The present SSSF does not include organic matter which is a very important part of the sea spray aerosol, in particular for the smallest particles (Facchini et al., 2008), which can act as cloud condensation nuclei. Therefore, we suggest to use the OSSA-SSSF to obtain the total sea spray aerosol fluxes and to derive the organic fraction by using the parameterization developed by Rinaldi et al. (2013) in a way it was used before in several other studies (O’Dowd et al., 2008; Vignati et al., 2010; Gantt et al., 2011). There is also evidence that micro-organisms affect the viscosity of sea water (Seuront et al., 2010) so that biological activity may be taken into account via the viscosity, like the effect of temperature and salinity; however, further studies and parameterizations are required on this topic in order to separate the different effects and relate the viscosity to observables like chlorophyll-a concentrations.

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